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(FILE 'HOME' ENTERED AT 19:58:24 ON 18 JUN 2008)

FILE 'REGISTRY' ENTERED AT 20:02:04 ON 18 JUN 2008

L1 STRUCTURE UPLOADED
L2 1 S L1 SSS
L3 27 S L1 SSS FULL

FILE 'CAPLUS' ENTERED AT 20:02:44 ON 18 JUN 2008

L4 1 S L3

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L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2008 ACS on STN
AN 2005:324121 CAPLUS

DN 142:392179

TI Preparation of acyloxydiphenylbutenylcinnamates as estrogen receptor modulator prodrugs.

IN Eaddy, John Fred, III; Heyer, Dennis; Katamreddy, Subba Reddy; Martin, Michael Tolar; McClure, Michael Scott; Randhawa, Amarjit Sab; Samano, Vicente; Ray, John Albert

PA Smithkline Beecham Corporation, USA

SO PCT Int. Appl., 78 pp.

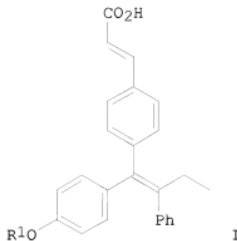
CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2005033056	A2	20050414	WO 2004-US32918	20041004
	WO 2005033056	A3	20050623		
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA,UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW:	BW, GH, GM, KE, LS, MW, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
EP	1692127	A2	20060823	EP 2004-809876	20041004
R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LT, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK, HR				
US	20070111971	A1	20070517	US 2006-575038	20060406
PRAI	US 2003-509678P	P	20031008		
	US 2003-514692P	P	20031027		
	WO 2004-US32918	W	20041004		
OS	CASREACT 142:392179; MARPAT 142:392179				
GI					



AB Title compds. (I; R1 = ACO, PO3H2; A = alkyl, aryl, heteroaryl, cycloalkyl, aminoalkyl, alkoxy, alkoxyalkyl, haloalkyl, heterocyclylalkyl), were prepared. Thus, I (R1 = H) (preparation given) and Et3N in THF at 5° were treated with propionyl chloride in THF followed by stirring for 1 h to give 64% I (R1 = EtCO). The latter orally in rats showed 86.6% bioavailability, vs. 5.7% for I (R1 = H).

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